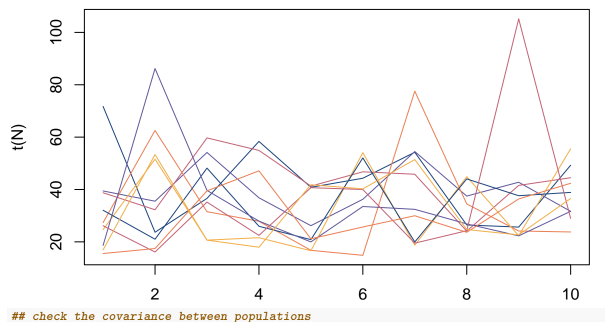
propagation_uncertainty_cov.R

katherine

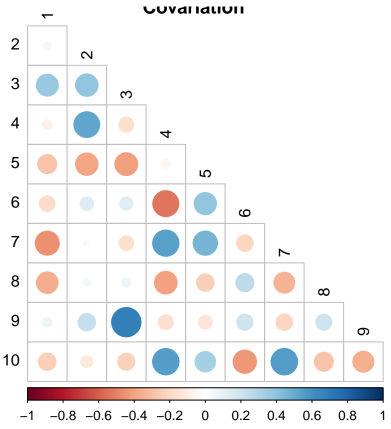
2022-06-29

```
# Demo the propagation of uncertainty in the calculation of the LPI
library(corrplot)
## simulate populations -----
# Function to simulate populations
\# For this demo, I am not adding any interactions between populations
sigma_process = 0.4
sigma_measure = 0.4
NO = 40
n_{pops} = 5
interaction_effect = 0.2
## carrying capacity scenarios ----
K_{increase} = NO + 5*c(0:10)
K_{stable} = rep(N0, 11)
K_{decline} = N0 - 2.5*c(0:10)
K = K_stable
lpiylim = c(0, 4)
sim_mech <- function(</pre>
    n_{pairs} = 10,
    timesteps = 10,
    NOi = NO, NOj = NO,
    lambda_i = 1.5, lambda_j = 1.5,
    alpha_ij = interaction_effect, alpha_ji = interaction_effect, # covariance
    process = proc,
    observation = obs,
    K){
  # set seed for randomisations
  set.seed(2)
  ## arguments: ## ----
  # npairs = number of pairs of populations to simulate
  # timesteps = number of time steps
```

```
# NOi = initial population size for set i
  # NOj = initial population size for set j
  # lambda i = true population growth rate for set i
  # lambda_j = true population growth rate for set j
  # alpha_ij = interaction coefficient (effect of set j on set i)
  # alpha_ji = interaction coefficient (effect of set i on set j)
  \# K = carrying \ capacity \ of \ the \ environment
  # process = absolute value of the to add to growth rates
  # observation = absolute value of the obs error limit to add to population sizes
  # add more time steps to allow lag
  timesteps = timesteps
  # initialize matrix to store results (population sizes)
  Ni <- as.matrix(rep(NOi, n_pairs))</pre>
  Nj <- as.matrix(rep(NOj, n_pairs))</pre>
  # initialize matrix to store growth rates
  dt_i <- matrix(NA, nrow = n_pairs, ncol = timesteps)</pre>
  dt_j <- matrix(NA, nrow = n_pairs, ncol = timesteps)</pre>
  \# Nt+1_i = Nt_i + rNt_i * ((1 - Nt_i/K_i) + alpha_ji*Nt_j/K_j)
  # calculate population sizes
  for(t in 1:timesteps){
    # generate growth rates from a lognormal distribution with process error
    r_i_error = rlnorm(n = n_pairs, meanlog = log(lambda_i), sdlog = process)
    r_j_error = rlnorm(n = n_pairs, meanlog = log(lambda_j), sdlog = process)
    # population i
    temp_i = Ni[,t]*(1 + r_i_error*(1 - (Ni[,t] + alpha_ij*Nj[,t])/K[t]))
    # if NAs or 0, set to 0.
    temp_i[which(is.na(temp_i))] <- 0</pre>
    temp i[which(temp i < 0)] <- 0</pre>
    Ni <- cbind(Ni, temp_i) # append resulting population size to results vector
    # population j
    temp_j = Nj[,t]*(1 + r_j = rror*(1 - (Nj[,t] + alpha_ji*Ni[,t])/K[t]))
    # if NAs or O, set to O.
    temp_j[which(is.na(temp_j))] <- 0</pre>
    temp_j[which(temp_j < 0)] <- 0</pre>
    Nj <- cbind(Nj, temp_j) # append resulting population size to results vector
    ## save generated growth rates
    dt_i[,t] <- r_i_error
    dt_j[,t] <- r_j_error
  # apply observation error on the calculated population sizes
  # pick number of lognormal distribution with a mean of N and an sd of observation error
  Ni <- apply(Ni, 1:2, function(x) {rlnorm(1, meanlog = log(x), sdlog = observation)})
```



cov(t(N)) %>% cov2cor() %>% corrplot(title = "Covariation", tl.col = "black", type = "lower", diag = FA

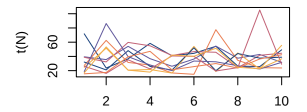


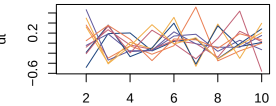
```
#### estimate population growth rate trend #### -----
# equation 2 ----
# expectation of the growth rate trend
eq2 <- function(N){
  d <- c()
  for(t in 2:length(N)){
    \label{eq:definition} d[t] \leftarrow \log 10(N[t]/N[t-1]) + (sigma_measure^2)/(2*(N[t-1]^2 - N[t]^2))
  }
  return(d)
}
dt <- apply(N, 1, eq2)</pre>
# equation 2
# just the growth rate part
eq2_donly <- function(N){
  eq2dt <- c()
  for(t in 2:length(N)){
    eq2dt[t] \leftarrow log10(N[t]/N[t-1])
  return(eq2dt)
}
donly <- apply(N, 1, eq2_donly)</pre>
# equation 2
```

```
# just the uncertainty part
eq2_varonly <- function(N){
  eq2var <- c()
  for(t in 2:length(N)){
    eq2var[t] <- (sigma_measure^2)/(2*((N[t-1])^2 - (N[t])^2))
    # should the population sizes be logged here?
    \#eg2var[t] \leftarrow (sigma_measure^2)/(2*(log(N[t-1])^2 - log(N[t])^2))
 }
 return(eq2var)
}
varonly <- apply(N, 1, eq2_varonly)</pre>
par(mfrow = c(2,2))
matplot(t(N), type = "l", main = "Population size", col = PNWColors::pnw_palette("Sunset2", 5), lty= 1)
matplot(dt, type = "l", main = "Growth rate (eq2)", col = PNWColors::pnw_palette("Sunset2", 5), lty= 1)
matplot(donly, type = "l", main = "Growth rate without uncertainty", col = PNWColors::pnw_palette("Suns
matplot(varonly, type = "1", main = "Measurement uncertainty", add = TRUE, col = PNWColors::pnw_palette
matplot(varonly, type = "1", main = "Measurement uncertainty", col = PNWColors::pnw_palette("Sunset2", ")
```

Population size

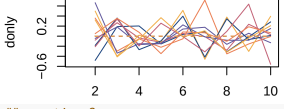
Growth rate (eq2)

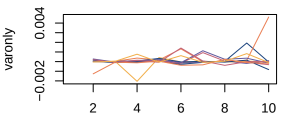




Growth rate without uncertainty

Measurement uncertainty

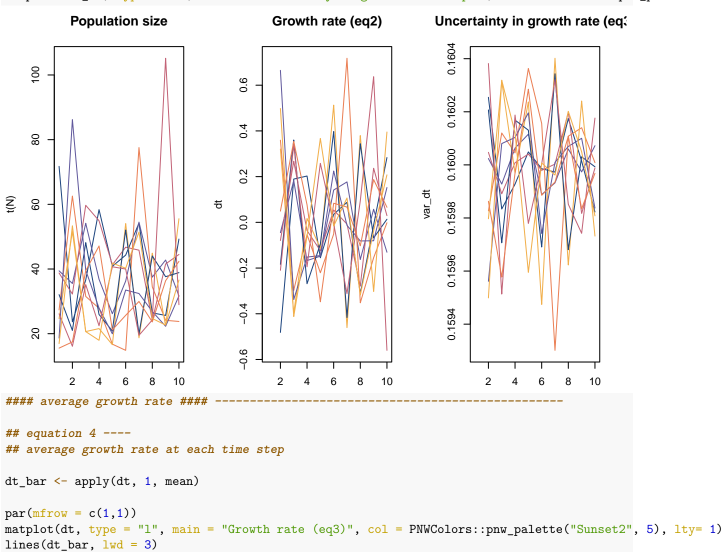




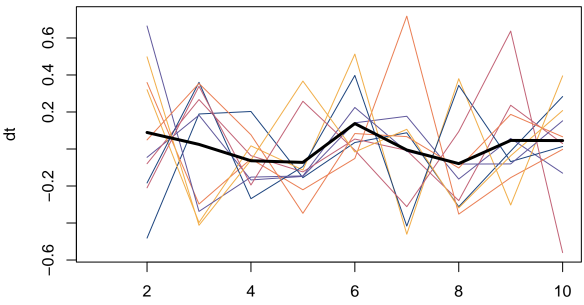
```
## equation 3 ----
eq3 <- function(N){
   eq3var <- c()
   for(t in 2:length(N)){
      eq3var[t] <- sigma_process^2 + (sigma_measure^2)*((1/((N[t])^2) - (1/(N[t-1])^2)))
   }
   return(eq3var)
}
var_dt <- apply(N, 1, eq3)

par(mfrow = c(1,3))</pre>
```

```
matplot(t(N), type = "l", main = "Population size", col = PNWColors::pnw_palette("Sunset2", 5), lty= 1)
matplot(dt, type = "l", main = "Growth rate (eq2)", col = PNWColors::pnw_palette("Sunset2", 5), lty= 1)
matplot(var_dt, type = "l", main = "Uncertainty in growth rate (eq3)", col = PNWColors::pnw_palette("Sunset2", 5)
```



Growth rate (eq3)

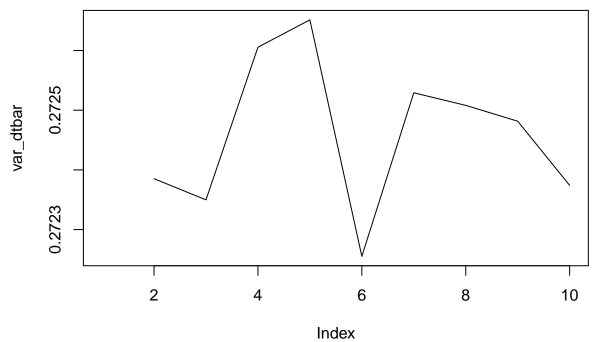


```
## equation 5 ----
## uncertainty in the average growth rate

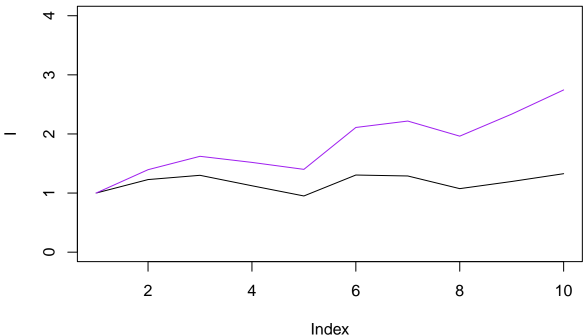
dt_cov <- cov(dt, use = "pairwise.complete.obs")
dt_cov <- dt_cov[which(lower.tri(dt_cov))]

var_dtbar = (1/n_pops)*(apply(var_dt, 1, sum) + 2*sum(dt_cov))

plot(var_dtbar, type = "l")</pre>
```



```
## equation 6 ----
## calculate LPI (without uncertainty correction)
# function to calculate LPI value without uncertainty correction
calclpi <- function(dt_bar){</pre>
  I = 1
  for(i in 2:length(dt_bar)){
    I[i] <- I[i-1]*10^dt_bar[i]</pre>
  }
  return(I)
}
## equation 7 ----
# function to calculate LPI value WITH uncertainty correction
calclpi_corrected <- function(dt_bar){</pre>
  I = 1
  for(i in 2:length(dt_bar)){
    I[i] <- I[i-1]*10^dt_bar[i] + 0.5*(10^dt_bar[i]*var_dtbar[i])</pre>
  return(I)
lpi_nocorrection = calclpi(dt_bar)
lpi_correction = calclpi_corrected(dt_bar)
plot(lpi_nocorrection, type = "l", ylab = "I", ylim = lpiylim)
lines(lpi_correction, col = "purple")
```



```
## equation 8 ----
# function to obtain the variance of the LPI
```

```
var_lpi <- (10^(2*dt_bar))*var_dtbar
plot(var_lpi, type = "l")</pre>
```

```
2 4 6 8 10
```

```
# render
# rmarkdown::render("propagation_uncertainty_cov.R", output_format = "pdf_document", output_file = "pro
# rmarkdown::render("propagation_uncertainty_cov.R", output_format = "pdf_document", output_file = "pro
# rmarkdown::render("propagation_uncertainty_cov.R", output_format = "pdf_document", output_file = "pro
```